

STIC Search Report Biotech-Chem Library

STIC Database Tracking Number: 114209

TO: Cybille Delacroix

Location: REM-4C70

Art Unit: 1614

February $\frac{18}{8}$, 2004

Case Serial Number: 09/700165

From: P. Sheppard

Location: Remsen Building

Phone: (571) 272-2529

sheppard@uspto.gov

Search Notes		
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SEARCH REQUEST FORM

Requestor's Delaunt	Serial 09 / 700 , 165
Date: $\frac{2-11-04}{}$ Pho	ne: <u>172-0572</u> Art Unit: <u>1414</u>
Search Topic: Please write a detailed statement of search topic. Describ that may have a special meaning. Give examples or relev a copy of the sequence. You may include a copy of the	be specifically as possible the subject matter to be searched. Define any terms vant citations, authors keywords, etc., if known. For sequences, please attach broadest and/or most relevant claim(s).
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Date completed: 0/18/04 Searcher: Sheppand Terminal time: Elapsed time:	Search Site Vendors STIC IG Suite CM-1 STN Pre-S Dialog
CPU time:	Type of Search APS N.A. Sequence Geninfo
Number of Searches: Number of Databases:	A.A. Sequence SDC Structure DARC/Questel

PTO-1500 (9-90)

USCOMM-DC 90-3952

delacroix 09_700165

=> FIL HCAPLUS
FILE 'HCAPLUS' ENTERED AT 14:46:40 ON 18 FEB 2004
USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.
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FILE COVERS 1907 - 18 Feb 2004 VOL 140 ISS 8 FILE LAST UPDATED: 17 Feb 2004 (20040217/ED)

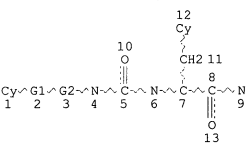
This file contains CAS Registry Numbers for easy and accurate substance identification.

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L3 STR



REP G1=(0-1) C REP G2=(0-1) CH2 NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 13

STEREO ATTRIBUTES: NONE

L5 1659 SEA FILE=REGISTRY SSS FUL L3

L6 STR

12 Cy 10	N~ G5 @14 15	H3C~~N~~CH3 16 @17 18
Cy G1 G2 - N - C - N - C - C - G4 1 2 3 4 5 6 7 9 0 13		

REP G1=(0-1) C REP G2=(0-1) CH2 REP G3=(0-3) CH2 VAR G4=NH/14

VAR G5=CY/OH/17/N/20/22

VAR G6=CH/28

VAR G7=CY/OH/N/20/22

NODE ATTRIBUTES:

DEFAULT MLEVEL IS ATOM

DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED

NUMBER OF NODES IS 29

STEREO ATTRIBUTES: NONE

L8 731 SEA FILE=REGISTRY SUB=L5 SSS FUL L6
L9 130 SEA FILE=HCAPLUS ABB=ON PLU=ON L8

L11 4 SEA FILE=HCAPLUS ABB=ON PLU=ON L9 AND (?SEXU? OR SEX? OR ?DYSFUN? OR ?ERECT? OR ?IMPOTEN? OR ?LIBIDO? OR ?ORGASM? OR

SUBSTITUTE OF SPRENDERS OF STREET, OF STREET, OF STREET, OF SPRENDERS

?VAGIN? OR ?DYSPAR?)

=> =>

=> D IBIB ABS HITRN L11 1-4

L11 ANSWER 1 OF 4 HCAPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER:

2003:58220 HCAPLUS 138:117676

DOCUMENT NUMBER: TITLE:

Linear and cyclic melanocortin receptor-specific

peptides, and therapeutic use

INVENTOR(S):

Sharma, Shubh D.; Shadiack, Annette M.; Yang, Wei;

Rajpurohit, Ramesh

PATENT ASSIGNEE(S):

Palatin Technologies, Inc., USA

PCT Int. Appl., 55 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent English

LANGUAGE:

SOURCE:

Englis

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2003006620	A2	20030123	WO 2002-US22196	20020711

delacroix 09 700165

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W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG

APPLN. INFO.:

| US 2001-304836P | P 20010711
      WO 2003006620
                            А3
                                    20031127
PRIORITY APPLN. INFO.:
                                                     US 2001-304836P P 20010711
                                 MARPAT 138:117676
OTHER SOURCE(S):
      Linear and cyclic peptides are provided which are specific to melanocortin
      receptors and which exhibit agonist, antagonist, or mixed
      agonist-antagonist activity. The peptides of the invention may be used to
      treat e.g. erectile dysfunction and eating disorders.
      488790-76-7
ΙT
      RL: PAC (Pharmacological activity); PRP (Properties); THU (Therapeutic
      use); BIOL (Biological study); USES (Uses)
           (linear and cyclic melanocortin receptor-specific peptides, and
          therapeutic use)
L11 ANSWER 2 OF 4 HCAPLUS COPYRIGHT 2004 ACS on STN
ACCESSION NUMBER:
                                 2002:869567 HCAPLUS
                                 137:370356
DOCUMENT NUMBER:
                                 Preparation and use of bombesin receptor antagonists
TITLE:
                                 for treatment of sexual dysfunction
                                 in males and females
                                 Gonzalez, Maria Isabel; Higginbottom, Michael; Stock,
INVENTOR(S):
                                 Herman Thijs; Pritchard, Martyn Clive; Pinnock, Robert
                                 Denham; Van der Graaf, Pieter Hadewijn; Naylor,
                                 Alisdair Mark; Wayman, Christopher Peter
PATENT ASSIGNEE(S):
                                 UK
                                 U.S. Pat. Appl. Publ., 105 pp., Cont.-in-part of U.S.
SOURCE:
                                 Pat. Appl. 2002 58,606.
                                 CODEN: USXXCO
DOCUMENT TYPE:
                                 Patent
                                 English
LANGUAGE:
FAMILY ACC. NUM. COUNT: 10
PATENT INFORMATION:
                                                        APPLICATION NO. DATE
      PATENT NO.
                           KIND DATE
                            ----
                                    _____
      US 2002169101
                            A1
                                    20021114
                                                        US 2001-999284 20011115
      US 2002058606 A1
                                                       US 2001-759777 20010112
                                    20020516
                                                    US 1999-133355P P 19990510
PRIORITY APPLN. INFO.:
                                                    WO 2000-GB1787 W 20000510
                                                    US 2000-700165 A2 20001109
                                                    US 2001-759777 A2 20010112
                                                    GB 2001-9910 A 20010423
                                                     GB 2001-11037 A 20010504
OTHER SOURCE(S):
                               MARPAT 137:370356
```

GI

AB Bombesin receptor antagonists have been found to be useful in the treatment of sexual dysfunction in both males and females. They may be selective BB1 antagonists or mixed BB1/BB2 antagonists. Combinations are disclosed of bombesin receptor antagonists with a range of other active compds., for example PDE5 inhibitors, NEP inhibitors and lasofoxifene. Prepn. of bombesin receptor antagonists consisting of .alpha.—Me tryptophane (e.g., I) or .alpha.—methylphenylalanine derivs. was given. In tests on sexually—dysfunctional male rats, it was concluded that I had a stimulatory effect, at the level of sexual desire, performance, and anorgasmy. In tests on sexually—dysfunctional female rats, it was concluded that I had a stimulatory effect on proceptivity, which was unaffected by repeated administration.

IT 428864-42-0

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (prepn. of as bombesin receptor antagonists for treatment of sexual dysfunction)

L11 ANSWER 3 OF 4 HCAPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER: 2002:391522 HCAPLUS

DOCUMENT NUMBER: 136:395983

TITLE: Bombesin receptor antagonists, and combinations with

other agents, for the treatment of sexual

dysfunction

INVENTOR(S): Gonzalez, Maria Isabel; Stock, Herman Thijs; Pinnock,

Robert Denham; Pritchard, Martyn Clive; Wayman, Christopher Peter; Van der Graaf, Pieter Hadewijn;

Naylor, Alisdair Mark; Higginbottom, Michael

PATENT ASSIGNEE(S): Warner-Lambert Company, USA

SOURCE: PCT Int. Appl., 225 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 10

PATENT INFORMATION:

PATENT NO.		KIND DATE			APPLICATION NO. DATE														
WO 2002040008		Α.	_				WO 2001-GB5018 20011114					1114							
WO 2002040008				-	2002														
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			CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	
			GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	ΚZ,	LC,	LK,	LR,	
			LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	ΜZ,	NO,	ΝZ,	PH,	PL,	
			PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	ТJ,	TM,	TR,	TT,	ΤZ,	UA,	UG,	
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BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
                                   20020523
      WO 2002040022
                            A1
                                                    WO 2000-GB4380 20001117
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RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
      AU 2002023802
                            Α5
                                   20020527
                                                     AU 2002-23802
                                                                          20011114
      EP 1333824
                            Α2
                                  20030813
                                                     EP 2001-994552
                                                                          20011114
               AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
                                   20030923
      BR 2001015364
                           Α
                                                     BR 2001-15364
                                                                          20011114
PRIORITY APPLN. INFO.:
                                                 WO 2000-GB4380
                                                                          20001117
                                                                      W
                                                 GB 2001-9910
                                                                      Α
                                                                          20010423
                                                 GB 2001-11037
                                                                      Α
                                                                          20010504
                                                 WO 2001-GB5018
                                                                      W
                                                                          20011114
OTHER SOURCE(S):
                               MARPAT 136:395983
      Bombesin receptor antagonists have been found to be useful in the
      treatment of sexual dysfunction in both males and
      females. They may be selective BB1 antagonists or mixed BB1/BB2
      antagonists. Combinations are disclosed of bombesin receptor antagonists
      with a range of other active compds., for example phosphodiesterase V
      inhibitors, neutral endopeptidase inhibitors, and lasofoxifene. Prepn. of
      compds. of the invention is described.
IT
      428864-42-0
      RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL
      (Biological study); USES (Uses)
          (bombesin receptor antagonists, and combinations with other agents, for
          treatment of sexual dysfunction)
L11 ANSWER 4 OF 4 HCAPLUS COPYRIGHT 2004 ACS on STN
ACCESSION NUMBER:
                               1995:812789 HCAPLUS
DOCUMENT NUMBER:
                               123:228906
                               Preparation of N-terminus modified analogs of
TITLE:
                               luteinizing hormone-releasing hormone (LHRH)
                               Haviv, Fortuna; Fitzpatrick, Timothy D.; Swenson, Rolf
INVENTOR(S):
                               E.; Nichols, Charles J.; Mort, Nicholas A.
                               Abbott Laboratories, USA
PATENT ASSIGNEE(S):
                               PCT Int. Appl., 42 pp.
SOURCE:
                               CODEN: PIXXD2
DOCUMENT TYPE:
                               Patent
                               English
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
      PATENT NO.
                                                    APPLICATION NO.
                           KIND
                                  DATE
                                                                          DATE
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      WO 9504540
                            A1
                                  19950216
                                                    WO 1994-US8577
                                                                          19940729
           W: CA, JP
           RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE
                                                     US 1993-103022
      US 5413990
                          A 19950509
                                                                          19930806
PRIORITY APPLN. INFO.:
                                                 US 1993-103022
                                                                          19930806
OTHER SOURCE(S):
                              MARPAT 123:228906
      N-acyldecapeptides X-A-B-C-D-E-F-G-H-I-J [I; X = acyl; A = D-Phe,
AB
      D-4-ClPhe, D-4-FPhe, D-3-(quinolin-3-yl)alanine, Sar, Gly, etc.; B =
      D-4-ClPhe, D-3,3-diphenylalanine, D-4-FPhe, D-3-(naphth-2-yl)alanine,
      D-Phe, D-3-(quinolin-3-yl)alanine; C = D-Ala, D-3-(benzo[b]thien-2-
      yl)alanine, Gly, D-3-(naphth-2-yl)alanine, D-3-(pyrid-3-yl)alanine,
      D-3-(quinolin-3-yl) alanine, D-3-(thiazol-2-yl) alanine; D=Gly, Ser,
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homo-Ser, Ser(CH2Ph), N.alpha.-C1-4 alkylserine; E = N.alpha.-R-3-R2-Ala, N.alpha.-R-3-R3-Lys, N.alpha.-R-Tyr, N.alpha.-R-Tyr(Me), N.alpha.-R-Phe, N.alpha.-cyclohexylalanine, N.alpha.-R-Gly, N.alpha.-R-Arg, N.alpha.-R1-His or -homo-His; wherein R = H, C1-4 alkyl; R2 = 4-(3-amino-1,2,4-triazol-5-yl)aminophenyl, 4-[[(3-amino-1,2,4-triazol-5yl)amino]methyl]phenyl, 4-(nicotinylamino)cyclohexyl, 4-nitrophenyl, 4-aminophenyl, etc.; R3 = N.epsilon.-nicotinyl or 3-amino-1,2,4-triazol-5yl; F = Gly, .beta.-alanine, D-citrulline, D-homo-citrulline, etc.; G = N.alpha.-R4-Leu, Gly, Sar, Pro, Val, N.alpha.-R4-L-cyclohexylalanine; R4 = H, C1-6 alkyl; H = L-citrulline, L-homo-citrulline, His, Lys(iso-Pr), N.alpha.-R4-Arg (wherein = same as above), homo-Arg, etc.; I = Pro, 4-hydroxy-L-proline, L-pipecoline, L-azetidine, N.alpha.-R4-Leu (wherein R4 = same as above), Sar, Gly, N.alpha.-R4-Ala, etc.; J = NHEt, N.alpha.-R4-D- or -L-Ala-NH2 (R4 = same as above), D-Ala-OH, D- or L-Glu, Sar-NH2, D-Ser-NH2, azaglycine, Gly-NH2] are prepd. These peptides I are potent antagonists of LHRH and are useful for suppressing the levels of sex hormones in mammals. Thus, CHO-D-2Nal-D-4-ClPhe-D-3Pal-Ser-MeTyr-D-Lys(Nic)-Leu-Lys(iso-Pr)-Pro-D-Ala-NH2 [2Nal = 3-(naphth-2-y1) alanine; 4-C1Phe = 3-(4-chloropheny1) alanine, 3Pal = 3-(pyrid-3-yl)alanine, Nic = nicotinyl] was prepd. by a Milligen-Biosearch 9,500 peptide synthesizer, involving sequential coupling of N-Boc-protected amino acids Boc-Pro-OH, Boc-Lys(Cbz,iso-Pr)-OH, Boc-Leu-OH, Boc-D-Lys(Nic)-OH, Boc-MeTyr(2,6-diCl-Bzl)-OH, Boc-Ser(Bzl)-OH, Boc-D-3Pal-OH, Boc-D-4ClPhe-OH, Boc-D-2Nal-OH and formic acid on a D-Ala-NH-resin (4-methylbenzhydrylamine resin). A total of 27 I were prepd. and in vitro showed pA2 values 8.8-11.46 in a test for LHRH antagonist potency, wherein the value of pA2 is the neg. logarithm of the concn. of the particular antagonist test compd. required to shift the response curve produced by the agonist leuprolide to two-fold higher concn. and typically pA2 values of .gtoreq.9.5 are indicative of good LHRH antagonist activity, with values of .gtoreq.10.0 being preferred. MeCH2CO-D-2Nal-D-4ClPhe-D-3Pal-Ser-MeTyr-D-Cit-Leu-Arg-Pro-D-Ala-NH2 (Cit = citrulline) showed the highest pA2 value (11.46).

IT 168158-00-7P 168158-04-1P 168158-06-3P 168158-08-5P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (prepn. of decapeptides as LH-releasing hormone (LHRH) antagonists)

=> =>

=> SELECT HIT RN L11 1-4 E541 THROUGH E546 ASSIGNED

=> FIL REG

FILE 'REGISTRY' ENTERED AT 14:47:04 ON 18 FEB 2004 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2004 American Chemical Society (ACS)

Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

STRUCTURE FILE UPDATES: 17 FEB 2004 HIGHEST RN 651291-85-9 DICTIONARY FILE UPDATES: 17 FEB 2004 HIGHEST RN 651291-85-9

TSCA INFORMATION NOW CURRENT THROUGH JULY 14, 2003

Please note that search-term pricing does apply when conducting SmartSELECT searches.

Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. For more information enter HELP PROP at an arrow prompt in the file or refer to the file summary sheet on the web at: http://www.cas.org/ONLINE/DBSS/registryss.html

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=> D HIS L12

(FILE 'HCAPLUS' ENTERED AT 14:44:08 ON 18 FEB 2004)

FILE 'HCAPLUS' ENTERED AT 14:46:40 ON 18 FEB 2004 SELECT HIT RN L11 1-4

FILE 'REGISTRY' ENTERED AT 14:47:04 ON 18 FEB 2004 L12 6 S E541-E546

=> =>

=> D IDE CAN L12 1-6

L12 ANSWER 1 OF 6 REGISTRY COPYRIGHT 2004 ACS on STN

RN 488790-76-7 REGISTRY

CN L-Tryptophanamide, 4-chloro-N-[[(2-phenylethyl)amino]carbonyl]-D-phenylalanyl-L-arginyl- (9CI) (CA INDEX NAME)

FS STEREOSEARCH

MF C35 H42 Cl N9 O4

SR CA

LC STN Files: CA, CAPLUS

Absolute stereochemistry.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1907 TO DATE)

1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 138:117676

L12 ANSWER 2 OF 6 REGISTRY COPYRIGHT 2004 ACS on STN

RN 428864-42-0 REGISTRY

CN 1H-Indole-3-propanamide, .alpha.-[[[[2,6-bis(dimethylamino)phenyl]amino]ca

rbonyl]amino]-N-(cyclohexylmethyl)-.alpha.-methyl- (9CI) (CA INDEX NAME)

MF C30 H42 N6 O2

SR CF

LC STN Files: CA, CAPLUS, USPATFULL

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

2 REFERENCES IN FILE CA (1907 TO DATE)

2 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 137:370356

REFERENCE 2: 136:395983

L12 ANSWER 3 OF 6 REGISTRY COPYRIGHT 2004 ACS on STN

RN 168158-08-5 REGISTRY

CN D-Alaninamide, 3-(2-naphthalenyl)-N-[[[2-(2-pyridinyl)ethyl]amino]carbonyl]-D-alanyl-4-chloro-D-phenylalanyl-3-(3-pyridinyl)-D-alanyl-L-seryl-N-methyl-L-tyrosyl-N5-(aminocarbonyl)-D-ornithyl-L-leucyl-L-arginyl-L-prolyl-, trifluoroacetate (salt) (9CI) (CA INDEX NAME)

FS PROTEIN SEQUENCE; STEREOSEARCH

MF C77 H100 Cl N19 O14 . x C2 H F3 O2

SR CA

LC STN Files: CA, CAPLUS

RELATED SEQUENCES AVAILABLE WITH SEQLINK

CM 1

CRN 168158-07-4

CMF C77 H100 C1 N19 O14

RELATED SEQUENCES AVAILABLE WITH SEQLINK

PAGE 1-A

PAGE 1-B

CM 2

CRN 76-05-1

CMF C2 H F3 O2

1 REFERENCES IN FILE CA (1907 TO DATE)

1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 123:228906

ANSWER 4 OF 6 REGISTRY COPYRIGHT 2004 ACS on STN L12

168158-06-3 REGISTRY RN

CN tyrosyl-N5-(aminocarbonyl)-D-ornithyl-L-leucyl-L-arginyl-L-prolyl-, trifluoroacetate (salt) (9CI) (CA INDEX NAME) PROTEIN SEQUENCE; STEREOSEARCH

FS

MF C76 H98 Cl N19 O14 . x C2 H F3 O2

SR CA

LC STN Files: CA, CAPLUS

RELATED SEQUENCES AVAILABLE WITH SEQLINK

CM

CRN 168158-05-2

C76 H98 Cl N19 O14 CMF

RELATED SEQUENCES AVAILABLE WITH SEQLINK

PAGE 1-B

CM 2

CRN 76-05-1 CMF C2 H F3 O2

1 REFERENCES IN FILE CA (1907 TO DATE)

1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 123:228906

L12 ANSWER 5 OF 6 REGISTRY COPYRIGHT 2004 ACS on STN

RN 168158-04-1 REGISTRY

CN D-Alaninamide, 3-(2-naphthalenyl)-N-[[(2-pyridinylmethyl)amino]carbonyl]-D-alanyl-4-chloro-D-phenylalanyl-3-(3-pyridinyl)-D-alanyl-L-seryl-N-methyl-L-tyrosyl-N5-(aminocarbonyl)-D-ornithyl-L-leucyl-L-arginyl-L-prolyl-, trifluoroacetate (salt) (9CI) (CA INDEX NAME)

FS PROTEIN SEQUENCE; STEREOSEARCH

MF C76 H98 Cl N19 O14 . x C2 H F3 O2

SR CA

LC STN Files: CA, CAPLUS

RELATED SEQUENCES AVAILABLE WITH SEQLINK

CM 1

CRN 168158-03-0

CMF C76 H98 Cl N19 O14

RELATED SEQUENCES AVAILABLE WITH SEQLINK

PAGE 1-B

$$(CH_2)_3$$

CM 2

CRN 76-05-1 CMF C2 H F3 O2

F-C-CO₂H

1 REFERENCES IN FILE CA (1907 TO DATE)
1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 123:228906

L12 ANSWER 6 OF 6 REGISTRY COPYRIGHT 2004 ACS on STN

RN 168158-00-7 REGISTRY

CN D-Alaninamide, N-[[[2-(4-morpholinyl)ethyl]amino]carbonyl]-3-(2-naphthalenyl)-D-alanyl-4-chloro-D-phenylalanyl-3-(3-pyridinyl)-D-alanyl-L-seryl-N-methyl-L-tyrosyl-N5-(aminocarbonyl)-D-ornithyl-L-leucyl-L-arginyl-L-prolyl-, trifluoroacetate (salt) (9CI) (CA INDEX NAME)

FS PROTEIN SEQUENCE; STEREOSEARCH

MF C76 H104 C1 N19 O15 . x C2 H F3 O2

SR CA

LC STN Files: CA, CAPLUS

RELATED SEQUENCES AVAILABLE WITH SEQLINK

CM 1

CRN 168157-99-1

CMF C76 H104 C1 N19 O15

RELATED SEQUENCES AVAILABLE WITH SEQLINK

PAGE 1-A

PAGE 1-B

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$$NH_3 \qquad NH$$

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$$NH_2 \qquad NH$$

$$NH_3 \qquad NH$$

$$NH_4 \qquad O \qquad NH$$

CM

CRN

76-05-1 C2 H F3 O2 CMF

1 REFERENCES IN FILE CA (1907 TO DATE)

1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 123:228906 => FIL HCAPLUS
FILE 'HCAPLUS' ENTERED AT 15:18:29 ON 18 FEB 2004
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PLEASE SEE "HELP USAGETERMS" FOR DETAILS.
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FILE COVERS 1907 - 18 Feb 2004 VOL 140 ISS 8 FILE LAST UPDATED: 17 Feb 2004 (20040217/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

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L38 ANSWER 1 OF 3 HCAPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER: 2002:391535 HCAPLUS

DOCUMENT NUMBER: 136:380143

TITLE: Treatment of sexual dysfunction

using bombesin antagonist

INVENTOR(S): Gonzalez, Maria Isabel; Higginbottom, Michael;

Pinnock, Robert Denham; Pritchard, Martyn Clive;

Stock, Herman Thijs

PATENT ASSIGNEE(S): Warner-Lambert Company, USA

SOURCE: PCT Int. Appl., 151 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 10 PATENT INFORMATION:

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      WO 2002040022
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RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
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      EP 1333829
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PRIORITY APPLN. INFO.:
                                                   GB 2001-9910
                                                                         A 20010423
                                                   GB 2001-11037
                                                                         A 20010504
                                                   WO 2001-GB5018
                                                                       W 20011114
      Bombesin receptor antagonists have been found to be useful in the
AB
      treatment of sexual dysfunction in both males and
      females. Prepn. of compds. of the invention is included.
      204067-01-6
TΤ
      RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL
      (Biological study); USES (Uses)
          (bombesin antagonists for treatment of sexual
          dysfunction)
                                        THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS
REFERENCE COUNT:
                                        RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L38 ANSWER 2 OF 3 HCAPLUS COPYRIGHT 2004 ACS on STN
ACCESSION NUMBER:
                                2002:368981 HCAPLUS
DOCUMENT NUMBER:
                                136:380137
TITLE:
                                Bombesin receptor antagonists, and preparation
                                thereof, for the treatment of sexual
                                dvsfunction
                                Gonzalez, Maria Isabel; Pinnock, Robert Denham;
INVENTOR(S):
                                Pritchard, Martyn Clive
PATENT ASSIGNEE(S):
                                UK
SOURCE:
                                U.S. Pat. Appl. Publ., 72 pp., Cont.-in-part of U.S.
                                Ser. No. 700,165.
                                CODEN: USXXCO
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delacroix 09 700165
DOCUMENT TYPE:
                         Patent
                         English
LANGUAGE:
FAMILY ACC. NUM. COUNT:
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PATENT INFORMATION:
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                      KIND DATE
                                            APPLICATION NO.
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     Bombesin receptor antagonists have been found to be useful in the
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     treatment of sexual dysfunction in both males and
     females.
IT
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     RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL
     (Biological study); USES (Uses)
        (bombesin receptor antagonists, prepn., and use for sexual
        dysfunction treatment, alone or with other agents)
L38 ANSWER 3 OF 3 HCAPLUS COPYRIGHT 2004 ACS on STN
                         2000:441789 HCAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                         133:74325
                         Preparation of non-peptide NK1 receptor antagonists
TITLE:
                         Creswell, Mark Wallace; Higginbottom, Michael;
INVENTOR(S):
                         Horwell, David Christopher; Lewthwaite, Russell
                         Andrew; Pritchard, Martyn Clive; Raphy, Jennifer
PATENT ASSIGNEE(S):
                         Warner-Lambert Company, USA
                         PCT Int. Appl., 131 pp.
SOURCE:
                         CODEN: PIXXD2
DOCUMENT TYPE:
                         Patent
                         English
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
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OTHER SOURCE(S): MARPAT 133:74325

AB Non-peptide acetamide derivs. R(CR1R2)m-X-CR3[(CH2)nR4]CONR9(CR5R7)qR6 [R = (un)substituted heterocyclyl, alkyl, hydroxy, alkoxy, halo,

PRIORITY APPLN. INFO.:

trifluoromethyl, carboxy, sulfonamido, nitro, 3.4-(H2NSO2)ClC6H3CH2, m-CF3SC6H4CH2, 3-(2-furyl)allyl, etc.; R1, R2, R3, R9 = H, alkyl; R4 =

US 2001-868449

US 1998-112725P P 19981218 WO 1999-US29592 W 19991214

A3 20010618

delacroix 09 700165

(un) substituted naphthyl or indolyl; R5, R7 = H, (CH2)pR10, where p = 1-3 and R10 = H, Me, CN, OH, OMe, CO2Me, NH2, NHMe, NMe2; R6 = (un) substituted Ph, naphthyl, heterocyclyl, alkyl, cycloalkyl, heterocycloalkyl, etc. or R5 and R6 form a bond; m = 0-3; n = 1, 2; q = 0-4; all stereoisomers possible] or their pharmaceutically acceptable salts were prepd. as specific NK1 antagonists. Thus, [R-(R*,S*)]-2-[(benzofuran-2-ylmethyl)amino]-3-(1H-indol-3-yl)-2-methyl-N-(1-phenylethyl)propionamide, prepd. by reductive condensation of 2-amino-3-(1H-indol-3-yl)-2-methyl-N-(1-phenylethyl)propionamide with 2-benzofurancarboxaldehyde using sodium triacetoxyborohydride, showed IC50 = 4.4 nM for in vitro human NK1 receptor binding.

IT 279222-39-8P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (prepn. of non-peptide NK1 receptor antagonists)

(prepn. of non-peptide NKI receptor antagonists)

REFERENCE COUNT: 10 THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

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=> FIL REG

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Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

STRUCTURE FILE UPDATES: 17 FEB 2004 HIGHEST RN 651291-85-9 DICTIONARY FILE UPDATES: 17 FEB 2004 HIGHEST RN 651291-85-9

TSCA INFORMATION NOW CURRENT THROUGH JULY 14, 2003

Please note that search-term pricing does apply when conducting SmartSELECT searches.

Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. For more information enter HELP PROP at an arrow prompt in the file or refer to the file summary sheet on the web at: http://www.cas.org/ONLINE/DBSS/registryss.html

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L39

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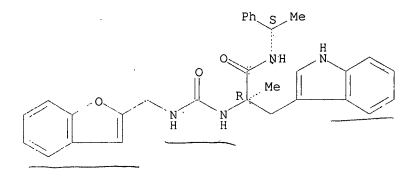
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- L39 ANSWER 1 OF 2 REGISTRY COPYRIGHT 2004 ACS on STN
- RN 279222-39-8 REGISTRY
- CN 1H-Indole-3-propanamide, .alpha.-[[((2-benzofuranylmethyl)amino]carbonyl]a mino]-.alpha.-methyl-N-[(1S)-1-phenylethyl]-, (.alpha.R)- (9CI) (CA INDEX NAME)
- FS STEREOSEARCH
- MF C30 H30 N4 O3

SR CA

LC STN Files: CA, CAPLUS, USPATFULL

Absolute stereochemistry. Rotation (+).



PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1907 TO DATE)

1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 133:74325

L39 ANSWER 2 OF 2 REGISTRY COPYRIGHT 2004 ACS on STN

RN 204067-01-6 REGISTRY

CN 1H-Indole-3-propanamide, N-[[1-(5-methoxy-2-pyridinyl)cyclohexyl]methyl].alpha.-methyl-.alpha.-[[[(4-nitrophenyl)amino]carbonyl]amino]-,
(.alpha.S)- (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 1H-Indole-3-propanamide, N-[[1-(5-methoxy-2-pyridinyl)cyclohexyl]methyl]-.alpha.-methyl-.alpha.-[[[(4-nitrophenyl)amino]carbonyl]amino]-, (S)-OTHER NAMES:

CN PD 176252

FS STEREOSEARCH

MF C32 H36 N6 O5

SR CA

LC STN Files: BIOSIS, CA, CAPLUS, SYNTHLINE, TOXCENTER, USPATFULL

 ${\tt Absolute \ stereochemistry.}$

9 REFERENCES IN FILE CA (1907 TO DATE)

9 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 140:70458

REFERENCE 2: 137:370356

REFERENCE 3: 137:41784

REFERENCE 4: 136:395983

REFERENCE 5: 136:380143

REFERENCE 6: 136:380137

REFERENCE 7: 134:172770

REFERENCE 8: 130:20201

REFERENCE 9: 128:205147

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=> D IDE CAN UREIDO

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=> D IDE CAN L33

L33 ANSWER 1 OF 1 REGISTRY COPYRIGHT 2004 ACS on STN RN 16518-43-7 REGISTRY CN Amidogen, (aminocarbonyl)- (9CI) (CA INDEX NAME) OTHER CA INDEX NAMES:

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Ureido (8CI)
CN
OTHER NAMES:
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LC
     STN Files:
       PIRA, PROMT, TOXCENTER
         (*File contains numerically searchable property data)
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7 REFERENCES IN FILE CA (1907 TO DATE)
7 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 137:169083

REFERENCE 2: 131:157549

REFERENCE 3: 127:206517

REFERENCE 4: 121:241460

REFERENCE 5: 80:89482

REFERENCE 6: 72:116747

REFERENCE 7: 67:16521

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FILE COVERS 1907 - 18 Feb 2004 VOL 140 ISS 8 FILE LAST UPDATED: 17 Feb 2004 (20040217/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

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GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 13

STEREO ATTRIBUTES: NONE

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RING(S) ARE ISOLATED OR EMBEDDED

NUMBER OF NODES IS 29

STEREO ATTRIBUTES: NONE

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L40	1	SEA FILE=HCAPLUS ABB=ON PLU=ON L26 NOT (L11 OR L38)

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L40 ANSWER 1 OF 1 HCAPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER:

2002:391709 HCAPLUS

DOCUMENT NUMBER:

136:386398

TITLE:

Preparation of bombesin receptor antagonists

INVENTOR(S):

Higginbottom, Michael; Pritchard, Martyn Clive; Stock,

Herman Thijs

PATENT ASSIGNEE(S):

Warner-Lambert Company, USA

SOURCE:

.PCT Int. Appl., 81 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PATENT NO. KIND DATE APPLICATION NO. DATE ----_____ WO 2001-EP14402 20011116 WO 2002040475 A1 20020523 W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG 20020522 GB 2000-28146 20001117 GB 2369118 A1 20020527 AU 2002-17095 AU 2002017095 Α5 20011116 20030813 EP 2001-996539 EP 1334102 A1 20011116 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR GB 2000-28146 A 20001117 PRIORITY APPLN. INFO.: WO 2001-EP14402 W 20011116 OTHER SOURCE(S): MARPAT 136:386398 Bombesin receptor antagonists (Ar)r-(CH2)j-(X)q-(CH2) kNR3CR5 (CH2Ar1) CONR4 (CH2) 1 (CR1R6) m (CH2) nR2 [j, n = 0-2; k, m, q, r = 0-2; k, m, q, q, r = 0-2; k, m, q,0 or 1; 1 = 0-3 (when r = 0, Ar is replaced by H); Ar = (un)substituted Ph, pyridyl, pyrimidyl, thienyl, furyl, imidazolyl, pyrrolyl or thiazolyl; Arl = any group for Ar or indolyl or pyridyl N-oxide; Rl = H, alkyl, (oxa, aza)cycloalkyl; R6 = H, Me or together with R6 forms a carbonyl group or a ring which can contain an oxygen or nitrogen atom; R3-R5 = H, alkyl; R2 = H, OH, alkoxy, NMe2, carbamoyl or certain ring structures; X is a divalent radical derived from isoxazole, pyridine, pyridazine, pyrimidine, etc.] or their pharmaceutically acceptable salts were prepd. The compds. of the invention have an affinity for the BB1 receptor and some of them also have affinity for the BB2 receptor. Accordingly they may be useful for the diagnosis, prevention, or treatment of male and female sexual dysfunction. Thus, (S)-3-(1H-indol)-3-yl)-N-[1-(5-methoxypyridin-2-yl)cyclohexylmethyl]-2-methyl-2-[4-(4-nitrophenyl)oxazol-2-ylamino] propionamide (1) was prepd. via amidation reaction and showed Ki = 4 or 24 nM in the BB1 and BB2 binding assay, resp. Compd. 1 was also assayed for female rat sexual proceptivity. THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 2 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT => SELECT HIT RN L40 1 NO E#s ASSIGNED => SELECT RN L40 1 E3 THROUGH E60 ASSIGNED

=> FIL REG

FILE 'REGISTRY' ENTERED AT 15:22:36 ON 18 FEB 2004 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2004 American Chemical Society (ACS)

Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

delacroix 09 700165

STRUCTURE FILE UPDATES: 17 FEB 2004 HIGHEST RN 651291-85-9 DICTIONARY FILE UPDATES: 17 FEB 2004 HIGHEST RN 651291-85-9

TSCA INFORMATION NOW CURRENT THROUGH JULY 14, 2003

Please note that search-term pricing does apply when conducting SmartSELECT searches.

Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. For more information enter HELP PROP at an arrow prompt in the file or refer to the file summary sheet on the web at: http://www.cas.org/ONLINE/DBSS/registryss.html

=> D HIS L41-

(FILE 'HCAPLUS' ENTERED AT 15:22:02 ON 18 FEB 2004) SELECT RN L40 1

FILE 'REGISTRY' ENTERED AT 15:22:36 ON 18 FEB 2004

L41 58 S E3-E60

L42 19 S L41 AND INDOL?

=> D IDE CAN L42 1-19

L42 ANSWER 1 OF 19 REGISTRY COPYRIGHT 2004 ACS on STN

RN 426267-06-3 REGISTRY

CN 1H-Indole-3-propanamide, .alpha.-methyl-N-[[1-(2-pyridinyl)cyclohexyl]methyl]-.alpha.-(4-quinolinylamino)-, (.alpha.S)-(9CI) (CA INDEX NAME)

FS STEREOSEARCH

MF C33 H35 N5 O

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL

Absolute stereochemistry.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

2 REFERENCES IN FILE CA (1907 TO DATE)

2 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 137:370356

REFERENCE 2: 136:386398

L42 ANSWER 2 OF 19 REGISTRY COPYRIGHT 2004 ACS on STN

RN 425639-33-4 REGISTRY

CN 1H-Indole-3-propanamide, .alpha.-methyl-.alpha.-[[(4-nitrophenyl)methyl]amino]-N-[[1-(2-pyridinyl)cyclohexyl]methyl]-, (.alpha.S)- (9CI) (CA INDEX NAME)

FS STEREOSEARCH

MF C31 H35 N5 O3

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL

Absolute stereochemistry.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

5 REFERENCES IN FILE CA (1907 TO DATE)

5 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 137:370356

REFERENCE 2: 136:395983

REFERENCE 3: 136:386398

REFERENCE 4: 136:380143

REFERENCE 5: 136:380137

L42 ANSWER 3 OF 19 REGISTRY COPYRIGHT 2004 ACS on STN

RN **425639-31-2** REGISTRY

CN 1H-Indole-3-propanamide, .alpha.-[(2-benzofuranylmethyl)amino].alpha.-methyl-N-[[1-(2-pyridinyl)cyclohexyl]methyl]-, (.alpha.S)(9CI) (CA INDEX NAME)

FS STEREOSEARCH

MF C33 H36 N4 O2

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL

5 REFERENCES IN FILE CA (1907 TO DATE)

5 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 137:370356

REFERENCE 2: 136:395983

REFERENCE 3: 136:386398

REFERENCE 4: 136:380143

REFERENCE 5: 136:380137

L42 ANSWER 4 OF 19 REGISTRY COPYRIGHT 2004 ACS on STN

RN **425639-28-7** REGISTRY

CN 1H-Indole-3-propanamide, .alpha.-methyl-.alpha.-[(2-phenylethyl)amino]-N-[[1-(2-pyridinyl)cyclohexyl]methyl]-, (.alpha.S)-(9CI) (CA INDEX NAME)

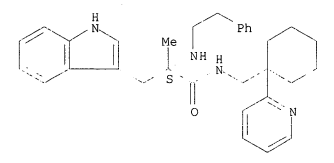
FS STEREOSEARCH

MF C32 H38 N4 O

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL

Absolute stereochemistry.



PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

5 REFERENCES IN FILE CA (1907 TO DATE) 5 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 137:370356

REFERENCE 2: 136:395983

REFERENCE 3: 136:386398

REFERENCE 4: 136:380143

REFERENCE 5: 136:380137

L42 ANSWER 5 OF 19 REGISTRY COPYRIGHT 2004 ACS on STN

RN **425639-22-1** REGISTRY

CN 1H-Indole-3-propanamide, .alpha.-methyl-.alpha.-[(6-phenyl-2-pyridinyl)amino]-N-[[1-(2-pyridinyl)cyclohexyl]methyl]-, (.alpha.S)-(9CI) (CA INDEX NAME)

FS STEREOSEARCH

MF C35 H37 N5 O

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL

Absolute stereochemistry.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

5 REFERENCES IN FILE CA (1907 TO DATE)

5 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 137:370356

REFERENCE 2: 136:395983

REFERENCE 3: 136:386398

REFERENCE 4: 136:380143

REFERENCE 5: 136:380137

L42 ANSWER 6 OF 19 REGISTRY COPYRIGHT 2004 ACS on STN

RN 425639-19-6 REGISTRY

CN 1H-Indole-3-propanamide, .alpha.-methyl-.alpha.-[(3-methylphenyl)amino]-N-[[1-(2-pyridinyl)cyclohexyl]methyl]-, (.alpha.S);

(9CI) (CA INDEX NAME)

FS STEREOSEARCH

MF C31 H36 N4 O

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL

Absolute stereochemistry.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

5 REFERENCES IN FILE CA (1907 TO DATE)

5 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 137:370356

REFERENCE 2: 136:395983

REFERENCE 3: 136:386398

REFERENCE 4: 136:380143

REFERENCE 5: 136:380137

L42 ANSWER 7 OF 19 REGISTRY COPYRIGHT 2004 ACS on STN

RN **425639-16-3** REGISTRY

CN 1H-Indole-3-propanamide, .alpha.-([1,1'-biphenyl]-2-ylamino)-.alpha.methyl-N-[[1-(2-pyridinyl)cyclohexyl]methyl]-, (.alpha.S)- (9CI) (CA
INDEX NAME)

FS STEREOSEARCH

MF C36 H38 N4 O

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL

5 REFERENCES IN FILE CA (1907 TO DATE)

5 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 137:370356

REFERENCE 2: 136:395983

REFERENCE 3: 136:386398

REFERENCE 4: 136:380143

REFERENCE 5: 136:380137

L42 ANSWER 8 OF 19 REGISTRY COPYRIGHT 2004 ACS on STN

RN **425639-13-0** REGISTRY

CN 1H-Indole-3-propanamide, .alpha.-methyl-N-[[1-(2-pyridinyl)cyclohexyl]methyl]-.alpha.-(5-pyrimidinylamino)-, (.alpha.S)-(9CI) (CA INDEX NAME)

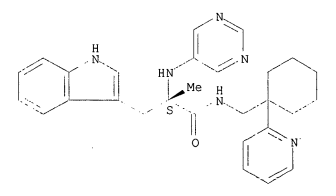
FS STEREOSEARCH

MF C28 H32 N6 O

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL

Absolute stereochemistry.



PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

delacroix 09 700165

5 REFERENCES IN FILE CA (1907 TO DATE) 5 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 137:370356

REFERENCE 2: 136:395983

REFERENCE 3: 136:386398

REFERENCE 4: 136:380143

REFERENCE 5: 136:380137

L42 ANSWER 9 OF 19 REGISTRY COPYRIGHT 2004 ACS on STN

RN 425639-07-2 REGISTRY

CN 1H-Indole-3-propanamide, .alpha.-methyl-.alpha.-(4-pyridinylamino)-N[[1-(2-pyridinyl)cyclohexyl]methyl]-, (.alpha.S)- (9CI) (CA INDEX NAME)

FS STEREOSEARCH

MF C29 H33 N5 O

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL

Absolute stereochemistry.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

5 REFERENCES IN FILE CA (1907 TO DATE) 5 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 137:370356

REFERENCE 2: 136:395983

REFERENCE 3: 136:386398

REFERENCE 4: 136:380143

REFERENCE 5: 136:380137

L42 ANSWER 10 OF 19 REGISTRY COPYRIGHT 2004 ACS on STN

RN 425639-04-9 REGISTRY

CN 1H-Indole-3-propanamide, .alpha.-(2-benzoxazolylamino)-.alpha.-methyl-N-[[1-(2-pyridinyl)cyclohexyl]methyl]-, (.alpha.S)- (9CI) (CA INDEX NAME)

FS STEREOSEARCH

MF C31 H33 N5 O2

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL

Absolute stereochemistry.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

5 REFERENCES IN FILE CA (1907 TO DATE)

5 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 137:370356

REFERENCE 2: 136:395983

REFERENCE 3: 136:386398

REFERENCE 4: 136:380143

REFERENCE 5: 136:380137

L42 ANSWER 11 OF 19 REGISTRY COPYRIGHT 2004 ACS on STN

RN 425639-02-7 REGISTRY

CN 1H-Indole-3-propanamide, N-[[1-(5-methoxy-2-pyridinyl)cyclohexyl]methyl]-.alpha.-methyl-.alpha.-[[4-(4-nitrophenyl)-2-thiazolyl]amino]-, (.alpha.S)- (9CI) (CA INDEX NAME)

FS STEREOSEARCH

MF C34 H36 N6 O4 S

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL

5 REFERENCES IN FILE CA (1907 TO DATE)

5 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 137:370356

REFERENCE 2: 136:395983

REFERENCE 3: 136:386398

REFERENCE 4: 136:380143

REFERENCE 5: 136:380137

L42 ANSWER 12 OF 19 REGISTRY COPYRIGHT 2004 ACS on STN

RN **425639-00-5** REGISTRY

CN 1H-Indole-3-propanamide, .alpha.-[(4-ethyl-2-oxazolyl)amino]-N-[[1-(5-methoxy-2-pyridinyl)cyclohexyl]methyl]-.alpha.-methyl-, (.alpha.S)(9CI) (CA INDEX NAME)

FS STEREOSEARCH

MF C30 H37 N5 O3

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL

5 REFERENCES IN FILE CA (1907 TO DATE)

5 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 137:370356

REFERENCE 2: 136:395983

REFERENCE 3: 136:386398

REFERENCE 4: 136:380143

REFERENCE 5: 136:380137

L42 ANSWER 13 OF 19 REGISTRY COPYRIGHT 2004 ACS on STN

RN 425638-98-8 REGISTRY

CN 1H-Indole-3-propanamide, N-[[1-(5-methoxy-2-pyridinyl)cyclohexyl]methyl]-.alpha.-methyl-.alpha.-[(4-phenyl-2-oxazolyl)amino]-, (.alpha.S)- (9CI) (CA INDEX NAME)

FS STEREOSEARCH

MF C34 H37 N5 O3

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL

5 REFERENCES IN FILE CA (1907 TO DATE)

5 REFERENCES IN FILE CAPLUS (1907 TO DATE)

1: 137:370356 REFERENCE

REFERENCE 2: 136:395983

REFERENCE 3: 136:386398

136:380143 REFERENCE 4:

REFERENCE 5: 136:380137

ANSWER 14 OF 19 REGISTRY COPYRIGHT 2004 ACS on STN L42

425638-96-6 REGISTRY RN

1H-Indole-3-propanamide, .alpha.-[[4-(4-cyanophenyl)-2-oxazolyl]amino]-N-[[1-(5-methoxy-2-pyridinyl)cyclohexyl]methyl]-.alpha.-CN methyl-, (.alpha.S)- (9CI) (CA INDEX NAME)
STEREOSEARCH

FS

C35 H36 N6 O3 MF

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL

5 REFERENCES IN FILE CA (1907 TO DATE)

5 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 137:370356

REFERENCE 2: 136:395983

REFERENCE 3: 136:386398

REFERENCE 4: 136:380143

REFERENCE 5: 136:380137

L42 ANSWER 15 OF 19 REGISTRY COPYRIGHT 2004 ACS on STN

RN **425638-92-2** REGISTRY

CN 1H-Indole-3-propanamide, .alpha.-methyl-.alpha.-[[4-(4-nitrophenyl)-2-coxazolyl]amino]-N-(2-oxo-2-phenylethyl)-, (.alpha.S)- (9CI) (CA INDEX NAME)

FS STEREOSEARCH

MF C29 H25 N5 O5

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL

5 REFERENCES IN FILE CA (1907 TO DATE)
5 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 137:370356

REFERENCE 2: 136:395983

REFERENCE 3: 136:386398

REFERENCE 4: 136:380143

REFERENCE 5: 136:380137

L42 ANSWER 16 OF 19 REGISTRY COPYRIGHT 2004 ACS on STN

RN 425638-90-0 REGISTRY

CN 1H-Indole-3-propanamide, N-[[1-(methoxymethyl)cyclohexyl]methyl].alpha.-methyl-.alpha.-[[4-(4-nitrophenyl)-2-oxazolyl]amino]-, (.alpha.S)(9CI) (CA INDEX NAME)

FS STEREOSEARCH

MF C30 H35 N5 O5

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL

Absolute stereochemistry.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

5 REFERENCES IN FILE CA (1907 TO DATE)

5 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 137:370356

REFERENCE 2: 136:395983

REFERENCE 3: 136:386398

REFERENCE 4: 136:380143

REFERENCE 5: 136:380137

L42 ANSWER 17 OF 19 REGISTRY COPYRIGHT 2004 ACS on STN

RN **425638-88-6** REGISTRY

CN 1H-Indole-3-propanamide, N-[[1-(5-methoxy-2-

pyridinyl)cyclohexyl]methyl]-.alpha.-methyl-.alpha.-[[4-(4-nitrophenyl)-2oxazolyl]amino]-, (.alpha.S)- (9CI) (CA INDEX NAME)

FS STEREOSEARCH

MF C34 H36 N6 O5

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL

Absolute stereochemistry.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

5 REFERENCES IN FILE CA (1907 TO DATE)

5 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 137:370356

REFERENCE 2: 136:395983

REFERENCE 3: 136:386398

REFERENCE 4: 136:380143

REFERENCE 5: 136:380137

L42 ANSWER 18 OF 19 REGISTRY COPYRIGHT 2004 ACS on STN

RN **204067-17-4** REGISTRY

CN 1H-Indole-3-propanamide, .alpha.-amino-.alpha.-methyl-N-[[1-(2-pyridinyl)cyclohexyl]methyl]-, (.alpha.S)- (9CI) (CA INDEX NAME) OTHER CA INDEX NAMES:

CN 1H-Indole-3-propanamide, .alpha.-amino-.alpha.-methyl-N-[[1-(2-pyridinyl)cyclohexyl]methyl]-, (S)-

FS STEREOSEARCH

MF C24 H30 N4 O

SR CA

LC STN Files: CA, CAPLUS, CASREACT, TOXCENTER, USPATFULL

7 REFERENCES IN FILE CA (1907 TO DATE)

7 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 137:370356

REFERENCE 2: 136:402022

REFERENCE 3: 136:395983

REFERENCE 4: 136:386398

REFERENCE 5: 136:380143

REFERENCE 6: 136:380137

REFERENCE 7: 128:205147

L42 ANSWER 19 OF 19 REGISTRY COPYRIGHT 2004 ACS on STN

RN **204067-16-3** REGISTRY

CN Carbamic acid, [(1S)-1-(1H-indol-3-ylmethyl)-1-methyl-2-oxo-2-[[[1-(2-pyridinyl)cyclohexyl]methyl]amino]ethyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN Carbamic acid, [1-(1H-indol-3-ylmethyl)-1-methyl-2-oxo-2-[[[1-(2-pyridinyl)cyclohexyl]methyl]amino]ethyl]-, 1,1-dimethylethyl ester, (S)-

FS STEREOSEARCH

MF C29 H38 N4 O3

SR CA

LC STN Files: CA, CAPLUS, CASREACT, TOXCENTER, USPATFULL

7 REFERENCES IN FILE CA (1907 TO DATE)
7 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 137:370356

REFERENCE 2: 136:402022

REFERENCE 3: 136:395983

REFERENCE 4: 136:386398

REFERENCE 5: 136:380143

REFERENCE 6: 136:380137

REFERENCE 7: 128:205147

Sheppard, Paula

From:

Moore, Peter

Sent:

Wednesday, February 18, 2004 3:06 PM

To: Cc:

Subject:

ABSS Notification

'dannie@cgen.com'; 'uspto-prj@cgen.com'

ABSS System Update (02/18/03)

The ABSS System Update:

<u>Note:</u> The databases on **ABSS01** will be updated once its job queues are cleared, **please hold all new requests to ABSS01** (#105136)

Reminder(s):

- (1) Database updates has completed on ABSS03h&p and ABSS06h&p.
- (2) When reporting an ABSS "<u>Server</u>" outage*, request that the Helpdesk create a "Critical Problem Notification (CPN)" for that server. All servers, related problems should also be "cc" to PTO's email "<u>System Software Division Unix</u>" (SSSD).

<u>Currently</u>, the following Compugen's production servers are up and running without any problems (
<u>ABSS03h</u>, <u>ABSS03p</u>, <u>ABSS04</u>, <u>ABSS05h</u>, <u>ABSS05p</u>, <u>ABSS06h</u> and <u>ABSS06p</u>). Please note additional instructions below:

RUSH Server(s):

> ABSS05h - Online, for "Rush" requests only!!! (Faster Server)

Servers open to all (No Rushes):

- > ABSS03h Online, can be loaded by everyone (Faster Server)
- > **ABSS03p Online,** can be loaded by everyone
- > ABSS04 Online, can be loaded by everyone (Faster Server)
- > ABSS05p Online, "Protein" requests only!!!
- ABSS06h Online, can be loaded by everyone to include "approved large search requests" (Faster Server)
- > **ABSS06p Online,** can be loaded by everyone

Additional Information:

- ABSS01: Online No new request 17Feb04 (2 BioXL/Hmm Jobs in Queue "119", Hours: "71:59")
- > ABSS03h: Online **Processing** 18Feb04 (2 BioXL/Hmm Jobs in Queue "13", Hours: "05:52")
- > ABSS03p: Online **Processing** 18Feb04 (4 BioXL/P Jobs in Queue "42", Hours: "18:05")
- > ABSS04: Online **Processing** 13Feb04 (2 BioXL/Hmm Jobs in Queue "**74**", Hours: "**77:04**")
- > ABSS05h: Online Processing 13Feb04 (2 BioXL/Hmm Jobs in Queue "96", Hours: "69:28")
- > ABSS05p: Online **Processing** 18Feb04 (4 BioXL/P Jobs in Queue "**79**", Hours: "**03:05**")
- > ABSS06h: Online **Processing** 17Feb04 (3 BioXL/Hmm Jobs in Queue "**104**", Hours: "**66:06**")
- > ABSS06p: Online **Processing** 18Feb04 (4 BioXL/P Jobs in Queue "**31**", Hours:

"19:24")

*Processing dates above reflect the day loaded into the ABSS system not the Examiner's submission date.

* Reserve for Testing (ABSS Testing team only):

- > ABSS02 Reserved Testing (2 BioXL/Hmm Jobs in Queue "0", Hours: "00:00")
- > Compugen1h Reserved Testing (1 BioXL/Hmm Jobs in Queue "0", Hours: "00:00")
- > Compugen1p Reserved Testing (2 BioXL/P Jobs in Queue "0", Hours: "00:00")

OFFLINE for Maintenance

> **ABSS01**: Staging for Dbase updates - **No new requests** (#105136)

Problem Reporting:

• Please continue to send email to the helpdesk, "CC" Walter and Peter for all servers, printers or workstations problems. This is a required procedure that must be followed to correctly login problem records.

For support when reporting a server, workstation or printer problem, please include the following.

- 1. Your name...
- 2. Automated Information System (AIS): ABSS
- 3. Name: (server, workstation or Printer name, i.e. ABSS06 or stic5 or QMS4)
- 4. Serial Number:
- 5. Location:
- 6. Description of problem:
- 7. Request an EA ticket number
- 8. Please **Call or E-Mail** the following:
 - If calling: 305-9000
 - If emailing: TO: HELPDESK9000

"CC" Rossy-Stiehl, Walter; Moore, Peter

Addition Notes:

All ABSS Servers Location: CPK2,11th floor - Main Computer Room, Suite 1100

Server Serial Numbers:

ABSS01......818FD15D ABSS02.....838F3699 ABSS03.....033H20FD ABSS04.....106A0914 ABSS05.....901H24F2

ABSS06......052H2C8D

Compugen1.....809FC101

Peter L. Moore III

System Software Division, CPK-2, Room C1009 Office: (703) 305-7795, Fax: (703)-308-4221

E-mail: peter.moore@uspto.gov <mailto:peter.moore@uspto.gov> <<mailto:peter.moore@uspto.gov>>

Sheppard, Paula

From:

Moore, Peter

Sent:

Wednesday, February 18, 2004_4:06 PM

To:

Wednesday, February 18, 2004 4:06 PM Alnaji, Read (Trawick); Arnold, Deirdre; Basker, Linda; Delaval, Jan; Hale, Mary; Hart, Edward; Hobbs, Lisa; Jarrell, Noble; Jones, Maude; O'Bryen, Barbara; Perkins, Leonard; Port, Toby; Rossy-Stiehl, Walter (Trawick); Ruhl, Mary Jane; Ruppel, Peggy; Schimmelbusch, Brian; Schreiber, David; Schulwitz, Paul; Shah, Arti; Shears, Beverly; Sheppard, Paula; Waclawiw, Alexandra; Westfall, Gary 'dannie@cgen.com'; 'uspto-prj@cgen.com'; Schimmelbusch, Brian; Hobbs, Lisa. ABSS System Update (02/18/03)-2

Cc: Subject:

Good afternoon,

Because of the large volumes of requests on ABSS04, we will need to stop accepting all new requests on ABSS04 starting 5pm tomorrow afternoon - Thursday, February 19, 2004, thanks.

Peter L. Moore III

System Software Division, CPK-2, Room C1009 Office: (703) 305-7795, Fax: (703)-308-4221

E-mail: peter.moore@uspto.gov"> << mailto:peter.moore@uspto.gov>